Corynebacterium marinum sp. nov. isolated from coastal sediment Zong-Jun Du, Elizabeth M. Jordan, Alejandro P. Rooney, Guan-Jun Chen and Brian Austin4

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1	Corynebacterium marinum sp. nov. isolated from coastal sediment in Qingdao,			
2	China			
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15	Running title: Corynebacterium marinum sp. nov.			
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17	The GenBank accession number for the 16S rRNA gene sequence of Corynebacterium marinum			
18	D7015 ^T is DQ219354. A representative phase-contrast micrograph of strain D7015 ^T and a table			
19	showing the cellular fatty acid profiles of strain D7015 ^T and related species are available as			
20	supplementary material in IJSEM Online.			

21	A taxonomic study was performed on strain D7015 ¹ , which was isolated from coastal sediment
22	close to a coal-fired power station in Qingdao, China. Strain D7015 ^T comprised Gram-positive,
23	non-motile diphtheroid rods, which grew in the presence of 0-8% (w/v) NaCl and at 4-37°C, with
24	optimum growth at 1% (w/v) NaCl and 30-32°C. The G+C content was 65.0 mol%. The major fatty
25	acids were $C_{18:1}$ ω 9c (56.18%), $C_{16:0}$ (38.02%), $C_{16:1}$ ω 7c (4.45%), $C_{18:0}$ (1.0%) and $C_{14:0}$ (0.35%).
26	On the basis of the morphological, physiological and phylogenetic characteristics, strain $D7015^{T}$
27	was classified in the genus Corynebacterium. It exhibited a 16S rRNA gene sequence similarity of
28	95.9% and a DNA:DNA relatedness value of 20.4% with Corynebacterium halotolerans DSM
29	44683 ^T . Strain D7015 ^T was sufficiently different from hitherto described <i>Corynebacterium</i> species
30	to be considered as a novel species. The name Corynebacterium marinum sp. nov. is proposed, with
31	strain $D7015^{T}$ (=CGMCC 1.6998 ^T =NRRL B-24779 ^T) as the type strain.

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The genus *Corynebacterium* was proposed by Lehmann & Neumann (1896) and represents a large
 group of Gram-positive, asporogenous, rod-shaped bacteria with high DNA G+C content.

Corynebacteria have been isolated from a wide range of environments, namely dairy products, soil, sewage, sediments, plant materials and aquatic sources, but the majority of novel species described in recent years have originated from human or animal clinical samples (e.g. Renaud *et al.*, 2007;

38 Yassin & Siering, 2008; Yassin, 2009; Funke et al., 2009). However, some corynebacterial species

39 have been isolated from the marine environment, and they may occur as part of the indigenous flora

40 of marine animals. For example, *Corynebacterium phocae* and *Corynebacterium caspium* were

isolated from seals (Pascual et al., 1998; Collins et al., 2004), Corynebacterium spheniscorum and

42 *Corynebacterium sphenisci* were recovered from wild penguins (Goyache *et al.*, 2003a, 2003b),

- 43 and Corynebacterium maris Coryn-1T was found in the mucus of the coral, Fungia granulose
- 44 (Ben-Dov et al., 2009). Here, we report the taxonomic characteristics of a novel Corynebacterium
- 45 species that originated from coastal sediment close to a coal-fired power station in Qingdao, China.

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47	Strain D7015 ^T was isolated from coastal sediment in 2000 (Du <i>et al.</i> , 2002). The isolate was grown
48	aerobically at 28°C on marine 2216E agar (MA; Difco) for 48 h. Cultures were maintained on MA
49	slants at room temperature, and stock cultures were kept in tryptone soya broth (Oxoid)
50	supplemented with 1% (w/v) NaCl (= TNB) and 20% (v/v) glycerol at -70°C. Identification of
51	strain D7015 ^T was performed as described by Jordan <i>et al.</i> (2007). For phenotypic tests, the strain
52	was grown on MA for 48 h at 28°C, and cells were resuspended in saline for use as an inoculum.
53	Tolerance of 1, 3, 5, 7 , 8 and 10% (w/v) NaCl was assessed on appropriately modified tryptone
54	soya agar (TSA; Oxoid). Growth in the absence of NaCl was assessed on plate count agar (PCA;
55	Oxoid). Inoculated plates were incubated at 28°C for up to 5 days. The effects of different
56	temperatures on growth were assessed on TSA plates supplemented with 1.0% (w/v) NaCl (TNA)
57	with incubation at 4, 10, 15, 28, 30, 32, 37, 42 and 45°C. The reduction of nitrate was assessed in
58	nitrate broth, prepared according to the method of Cowan & Steel (1974), and incubated at room
59	temperature for 10 days. Oxidase and catalase activities were determined by using standard
60	methods. The culture was characterized biochemically using the API Coryne, API 50CH and API
61	ZYM systems according to the manufacturer's instructions (bioMérieux). Measuring turbidity was
62	used to evaluate the growth of strain $D7015^{T}$ in the API 50CH and the API 20NE systems. The API
63	50 CH strips were read after 7 days incubation at 28°C.
64	

The almost-complete 16S rRNA gene sequence (1444 nt) of strain D7015^T was obtained using the universal primers 27f and 1492r (MWG Biotech; Lane, 1991). The 16S rRNA gene sequence of strain D7015^T was submitted to GenBank and EMBL to search for similar sequences using the BLAST algorithm. A phylogenetic dendrogram of strain D7015^T and some closely related members of the genus *Corynebacterium* based on 16S rRNA gene homology was constructed using the neighbor-joining method of the MEGA software version 4.1 (Tamura *et al.*, 2007). The

resultant tree topologies were evaluated by bootstrap analysis based on 1000 replicates.

73	Cellular fatty acids were determined on a 3 day old culture grown on marine 2216E agar plates after
74	incubation at 28°C. The fatty acids were extracted, methylated and analyzed using the standard
75	MIDI (Microbial Identification) system (Sasser, 1990). The G+C content of the DNA was
76	determined directly by high pressure liquid chromatography according to a method described
77	previously (Tamaoka & Komagata, 1984; Mesbah et al., 1989). DNA:DNA hybridization between
78	strain D7015 ^T and <i>Corynebacterium halotolerans</i> DSM 44683 ^T was carried out by applying the
79	optical renaturation method (De Ley et al., 1970; Huss et al., 1983; Jahnke, 1992) under optimal
80	hybridization conditions.
81	
82	Microscopic examination of strain D7015 ^T suggested that it belonged to the genus
83	Corynebacterium, as cells stained Gram-positive and produced short, diphtheroid rods with some
84	of the cells arranged in a V formation due to their snapping division (Collins & Cummins, 1986). A
85	representative phase-contrast micrograph of strain D7015 ^T is available as supplementary material
86	in IJSEM Online. Strain D7015 ^T was non-motile and non-endospore-forming. The isolate was
87	catalase-positive and oxidase-negative. The following results of carbon source assimilation were
88	positive in API 50CH: aesculin ferric citrate, salicin, D-maltose and glycogen. Strain D7015 ^T
89	displayed a numerical profile of 3200127 with the commercial API Coryne system, which
90	corresponded to a "doubtful" identification as Corynebacterium glucuronolyticum (with a
91	confidence level of 99.2%). However, further biochemical analyses, using API ZYM and
92	additional phenotypic tests revealed that strain D7015 ^T could be distinguished from C.
93	glucuronolyticum on the basis of its ability to produce α -chymotrypsin and its inability to produce
94	esterase lipase (C4). Strain D7015 ^T produced acid from glucose, maltose, sucrose and glycogen,
95	but not from ribose, xylose, mannitol and lactose. Strain $D7015^{T}$ was also different from <i>C</i> .

spheniscorum notably in the inability of the latter to reduce nitrate, or produce β-glucuronidase, αchymotrypsin or naphthol-AS-BI-phosphohydrolase. The complete morphological and biochemical characteristics for strain D7015^T are given in the species description.

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Phylogenetic analyses performed with nearly complete sequences of members of closely related 100 species (Fig. 1) showed that no sequence available in the GenBank database exhibited more than 101 102 96% similarity. The sequence similarity observed between the new isolate and its closest relative, C. halotolerans, was 95.9%, which is a lower value than the borderline used for defining bacterial 103 species (i.e. 97%) as proposed by Stackebrandt & Goebel (1994). C. halotolerans was isolated 104 from a saline soil sample in China (Chen *et al.*, 2004). For strain D7015^T, growth was not observed 105 106 at 10% (w/v) NaCl. However, optimum growth of C. halotolerans was in 10% (w/v) KCl, NaCl or MgCl₂·6H₂O. These two strains were also different biochemically, with strain D7015^T able to 107 hydrolyse starch and Tween 20-80, but unable to produce esterase lipase (C4). The DNA:DNA 108 109 relatedness value of $20.4 \pm 0.1\%$ (experiment repeated twice) was significantly lower than 70%, which is considered to be the threshold value for the delineation of genomic species (Wayne et al., 110 1987). Clearly, strain D7015^T differed biochemically from all of its closest relatives, both in its 111 ability to hydrolyse casein and in its inability to produce esterase lipase C4. The phenotypic 112 features that differentiate strain D7015^T from its closest phylogenetic relatives are provided in 113 114 Table 1.

115

116 The cellular fatty acids in strain D7015^T were $C_{18:1}\omega9c$ (56.18%), $C_{16:0}$ (38.02%), $C_{16:1}\omega7c$ (4.45%), 117 $C_{18:0}$ (1.0%) and $C_{14:0}$ (0.35%). The genomic DNA G+C content was 65.0 mol%.

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Based on these molecular, chemotaxonomic and phenotypic results, it is proposed that strain $D7015^{T}$ should be classified as a new species of the genus *Corynebacterium*, for which the name

121 *Corynebacterium marinum* sp. nov. is proposed.

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123 Description of *Corynebacterium marinum* sp. nov.

124 *Corynebacterium marinum* (ma.ri'num. L. neut. adj. *marinum* of the sea, marine)

125

126 Cells are short Gram-positive, non-motile, diphtheroid rods; some of the cells are arranged in a V formation. Colonies on marine 2216E agar medium are circular, erose, convex, yellow and of a 127 creamy consistency and are 0.5-1.5 mm in diameter after 48 h at 28°C. Facultatively anaerobic, 128 129 catalase-positive and oxidase-negative. Bacteria are methyl-red negative, Voges-Proskauer positive, reduce nitrate, and lyse horse blood cells. Aesculin and urea are not hydrolysed, but casein 130 131 is digested and starch and pullulan are hydrolysed. Gelatin is not liquefied. Tween 20-80 are 132 hydrolysed. Cells grow in the presence of 0-8% (w/v) NaCl and at 4-37°C. Prolific growth occurs 133 at 30-32°C in media that contain 1% (w/v) NaCl. Using the API 50CH system, aesculin ferric 134 citrate, salicin, D-maltose, and glycogen are utilized. Acid is produced from glucose, maltose, 135 sucrose and glycogen, but not from ribose, xylose, mannitol or lactose. Activities for esterase lipase (C8), leucine arylamidase, α -chymotrypsin, naphthol-as-bi-phosphohydrolase, pyrazinamidase and 136 β -glucuronidase are positive. Esterase lipase (C4), lipase (C14), valine arylamidase, cystine 137 arylamidase, trypsin, alkaline phosphatase, pyrrolidonyl arylamidase, acid phosphatase, 138 α -galactosidase, β -galactosidase, α -glucosidase, β -glucosidase, N-acetyl- β -glucosaminidase, 139 140 α -mannosidase and α -fucosidase are negative. The type strain is resistant to nalidixic acid (30 µg), nitrofurantoin (50 μ g), sulphamethizole (200 μ g), tetracycline (100 μ g) and cotrimoxazole (25 μ g), 141 but sensitive to ampicillin (25 µg), chloramphenicol (50 µg), gentamycin (10 µg), kanamycin (30 142 143 μ g), carbenicillin (100 μ g) and streptomycin (25 μ g) as determined by antibiotic discs. Major fatty acids produced are $C_{18:1}$ ω 9c (56.18%), $C_{16:0}$ (38.02%), $C_{16:1}$ ω 7c (4.45%), $C_{18:0}$ (1.0%) and $C_{14:0}$ 144 (0.35%). The percentage whole-cell fatty acid compositions of strain D7015^T and related species is 145

available as supplementary material in IJSEM Online. The genomic DNA G+C content is 65.0
mol% for the type strain.

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149 The type strain is $D7015^{T}$ (=CGMCC 1.6998^T = NRRL B-24779^T), which was isolated from coastal 150 sediment close to a coal-fired power station in Qingdao, China.

151

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- **Table 1.** Comparison of strain $D7015^{T}$ with the phylogenetically related species of the genus
- 230 Corynebacterium
- 231 Species: 1, Strain D7015^T, 2, *Corynebacterium efficiens* DSM 44549^T (Fudou *et al.*, 2002); 3,
- *Corynebacterium halotolerans* DSM 44683^T (Chen *et al.*, 2004). +, positive; -, negative; ND, no

Characteristic	1	2	3
Starch hydrolysis	+	ND	-
Esterase lipase (C4)	-	ND	+
Growth at 45 °C	-	+	ND
Growth in 10% (w/v) NaCl	-	+	+
Acid produced from:			
ribose	-	+	-
maltose	+	+	-
glycogen	+	-	-
DNA G+C content (mol%)	65.0	59.0-60.2	63.0

233 data available.

245 Figure legend:

- Fig. 1. Neighbor-joining phylogeny of *Corynebacterium marinum* and closely related species.
- 247 Numbers along branches represent bootstrap values; only values greater than 50% are shown.
- 248 GenBank accession numbers for the 16S rRNA sequences used in the tree reconstruction are given
- in the parentheses next to each taxon. Bar, 0.01 substitutions per nucleotide.

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